

Correction: Dynamic behavior of rearranging carbocations – implications for terpene biosynthesis

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Correction

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This correction refers to *Beilstein J. Org. Chem.* **2016**, *12*, 377–390. doi:10.3762/bjoc.12.41

The originally published Figure 6 had several mis-drawn structures with charges in incorrect locations. These errors have been corrected in the new version (see Figure 1).

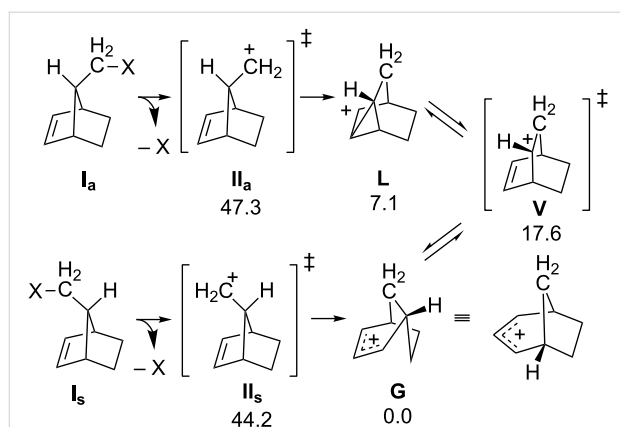


Figure 1: Corrected Figure 6 of the original article. The portion of the norborn-2-en-7-ylmethyl cation PES examined by Ghigo et al. [60]. Energies reported are electronic energies, including zero-point corrections (ZPE), at the B3LYP/6-31G(d) level of theory and are all relative to that of **G** [61–63] (for references see original article).

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